

(19)



Europäisches Patentamt

European Patent Office

Office européen des brevets



(11)

EP 0 926 165 A1

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication:

30.06.1999 Bulletin 1999/26

(51) Int. Cl.<sup>6</sup>: C08F 4/654, C08F 10/00

(21) Application number: 98660147.4

(22) Date of filing: 21.12.1998

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU  
MC NL PT SE

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 23.12.1997 FI 974621

23.12.1997 FI 974622

(71) Applicant: BOREALIS A/S

2800 Lyngby (DK)

(72) Inventors:

- Garoff, Thomas  
00840 Helsinki (FI)
- Leinonen, Timo  
06750 Tolkkinen (FI)
- Ala-Huikku, Sirpa  
00970 Helsinki (FI)

(74) Representative:

Träskman, Berndt Hilding  
Berggren Oy Ab,  
P.O. Box 16  
00101 Helsinki (FI)

(54) **Catalyst component comprising magnesium, titanium, a halogen and a electron donor, its preparation and use**

(57) The invention relates to a novel process for the preparation of an olefin polymerization catalyst component comprising magnesium, titanium, a halogen and an electron donor. The process comprises the steps of:

electron donor by replacement of its halogen by said alkoxy moiety.

(i) reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxy linked to magnesium, a complex containing a magnesium dihalide and an alcohol, and a non-complex magnesium dialkoxide, with a halogen compound (b) being capable of forming the electron donor by replacement of its halogen by said alkoxy moiety, to give an intermediate (ab), and  
(ii) reacting said intermediate (ab) with a titanium halide (c), or

(i)' reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxy linked to magnesium, and a complex containing a magnesium dihalide and an alcohol, with a titanium halide (c), to give an intermediate (ac), and  
(ii)' reacting said intermediate (ac) with a halogen compound (b) being capable of forming the

EP 0 926 165 A1

**Description**

[0001] The invention relates to a process for the preparation of an olefin polymerization catalyst component containing magnesium, titanium, halogen and an electron donor. The invention also relates to such a catalyst component and its use for the polymerization of  $\alpha$ -olefins such as propene.

**Background of the invention**

[0002] Generally, so called Ziegler-Natta catalyst components of the above kind have been prepared by reacting a magnesium halide-alcohol complex with a titanium tetrahalide and an electron donor which usually is a phthalic acid diester. The preparation involves the use of large amounts of reagents and washing liquids, which are difficult to handle. Additionally, byproducts are formed, which cannot easily be regenerated or destroyed, but form an environmental problem.

[0003] For example, the preparation of a conventional polypropene catalyst component involves the reaction of a magnesium dichloride-alcohol complex with titanium tetrachloride to give reactive  $\beta$ -magnesium dichloride as intermediate and hydrogen chloride and titanium alkoxy trichloride as byproducts. Then, the reactive  $\beta$ -magnesium dichloride intermediate is activated with further titanium tetrachloride to give said catalyst component (the treatment with a titanium halide such as titanium tetrachloride is henceforth called titanation).

[0004] The titanium alkoxy trichloride byproduct formed in the titanation is a catalyst poison and must be carefully removed by extensive washing using large amounts of titanium tetrachloride. Further, the titanium alkoxy trichloride must be carefully separated from the titanium tetrachloride washing liquid, if the latter is to be reused e.g. for activating the reactive  $\beta$ -magnesium dichloride. Finally, the titanium alkoxy trichloride is a problem waste, which is difficult to dispose of.

[0005] Thus, in a typical propene polymerization catalyst component preparation involving two titanations and three heptane washes, one mol of produced catalyst component (mol Mg) requires about 40 mol of titanium tetrachloride e.g. as washing liquid to be circulated (see Table 15 below), and produces as problem waste about three mol of titanium alkoxy trichloride as well as about three mol of hydrogen chloride.

[0006] Sumitomo, EP 0 748 820 A1 (hereinafter referred to as "Sumitomo"), has prepared dialkoxy magnesium, reacted it with titanium tetrachloride to form an intermediate and then reacted the intermediate with phthalic acid dichloride to form a catalytically active propene polymerization catalyst component. The activity was raised by repeated titanations, as well as repeated washes with toluene and hexane. See page 10, lines 14 to 37, of said publication.

[0007] Said process of Sumitomo has avoided the reaction between the magnesium dichloride-alcohol complex and titanium tetrachloride, and thereby eliminated the formation of catalytically poisonous titanium alkoxy trichloride byproduct. However, as much as four titanations and hydrocarbon treatments are still needed to give satisfactory catalytic activity.

**Description of the Invention**

[0008] The purpose of the present invention is to provide a process which results in a catalyst component having satisfactory activity without producing harmful byproducts such as said titanium alkoxy trichloride or requiring the use of a large amounts of titanation reagent and/or washing liquid.

[0009] The problem described above has now been solved with a novel process for the preparation of a catalyst component of the above type, which is mainly characterized by the steps of:

[45] (i) reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxide linked to magnesium, a complex containing a magnesium dihalide and an alcohol, and a non-complex magnesium dialkoxide, with a halogen compound (b) being capable of forming the electron donor by replacement of its halogen by said alkoxy moiety, to give an intermediate (ab), and

[50] (ii) reacting said intermediate (ab) with a titanium halide (c), or

[55] (i)' reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxide linked to magnesium, and a complex containing a magnesium dihalide and an alcohol, with a titanium halide (c), to give an intermediate (ac), and

(ii)' reacting said intermediate (ac) with a halogen compound (b) being capable of forming the electron donor by replacement of its halogen by said alkoxy moiety.

[0010] It was found by the applicant, that the activity of a stoichiometric catalyst component, comprising a magnesium dihalide, a titanium tetrahalide and an electron donor, is the higher, the more magnesium dihalide it contains. Thus it is believed, that the purpose of the repeated toluene washes of e.g. Sumitomo has partly been to remove titanium tetrachloride and electron donor from the catalyst component precursor in order to raise the magnesium dichloride content and thus the catalytic activity, of the final catalyst component. The present invention solves the problem in another way. In the claimed process, magnesium dihalide is included or synthesized as part of the reacting material before any titanation takes place, and thus, the need for repeated cycles of titanation and washing is significantly reduced.

[0011] Preferably one, most preferably all of steps (i), (ii), (i)' and (ii)' are performed in solution. Then, the reaction product of step (ii) or (ii)' is preferably recovered by precipitation.

[0012] According to one embodiment of the present invention, said compounds (a), (b) and (c) are in the claimed process contacted in essentially stoichiometric amounts. According to another embodiment, a stoichiometric excess, preferably a 5-20 fold excess, of said titanium halide (c) with respect to said magnesium compound (a), gives even better results.

[0013] Said halogen compound (b) used in the claimed process is an electron donor precursor, i.e. itself capable of forming the electron donor of the catalyst component by replacement of its halogen by an alkoxy group. By electron donor is in this connection meant an electron donor which forms a part of the titanous catalyst component produced by the claimed process and is in the art also called an internal electron donor. Such halogen compounds (b) are, e.g., C<sub>1</sub>-C<sub>20</sub> alkyl halides, C<sub>7</sub>-C<sub>27</sub> aralkyl halides and C<sub>2</sub>-C<sub>22</sub> acyl halides, which react with alkoxy compounds to replace their halogen with the alkoxy group of the alkoxy compound and form e.g. the corresponding ethers and esters acting as internal electron donors.

[0014] Preferably, said halogen compound (b) is an organic acid halide having the formula R"(COX)<sub>n</sub>, wherein R" is an n-valent organic group having 1-20 carbon atoms, preferably an n-valent benzene ring, X is a halogen, preferably chlorine, and n is the valence of R" and is an integer 1 to 6, preferably 1, 2, 3 or 4, more preferably 2. Most preferably, said halogen compound is phthalic acid dichloride Ph(COCl)<sub>2</sub>, wherein Ph is o-phenylene. Correspondingly, the electron donor formed therefrom is preferably an organic acid ester having the formula R"(COOR)<sub>n</sub>, wherein R is an n-valent C<sub>1</sub>-C<sub>20</sub> aliphatic group or an n-valent C<sub>7</sub>-C<sub>27</sub> araliphatic group and R" and n are the same as above, and more preferably a phthalic acid diester Ph(COOR)<sub>2</sub>, wherein R is a C<sub>1</sub>-C<sub>20</sub> alkyl or a C<sub>7</sub>-C<sub>27</sub> aralkyl, more preferably a C<sub>1</sub>-C<sub>16</sub> alkyl. Most preferably said electron donor is diethyl phthalate.

[0015] The titanium halide (c) used in the claimed process is preferably a titanium halide of the formula (OR')<sub>k</sub>TiX<sub>4-k</sub>, wherein R' is an alkyl group having 1 to 10 carbon atoms or an aralkyl group having 7 to 16 carbon atoms, X is a halogen and k is 0 to 3. More preferably, said titanium halide (c) is a titanium tetrahalide TiX<sub>4</sub>, wherein X is the same as above, most preferably titanium tetrachloride TiCl<sub>4</sub>.

[0016] It is preferable, if said titaniumless magnesium compound (a) is not a part of a solid magnesium halide, e.g. in the form of complex molecules on the surface of a solid magnesium halide carrier, but form a separate compound with an essentially stoichiometric composition. Often, said titaniumless magnesium compound is a complex. A complex is, according to Römpps Chemie-Lexicon, 7. Edition, Franckh'sche Verlagshandlung, W. Keller & Co., Stuttgart, 1973, page 1831, "a derived name of compounds of higher order, which originate from the combination of molecules, - unlike compounds of first order, in the creation of which atoms participate".

[0017] According to one embodiment of the invention, the titaniumless magnesium compound (a) used in the claimed process is a titaniumless complex of the formula [MgX<sub>2</sub>]<sub>x</sub> · [K(OR)<sub>m</sub>]<sub>y</sub>, wherein X is a halogen, K is hydrogen, a metal of group 1, 2 or 13 of the Periodic Table, R is an alkyl having 1 to 20 carbon atoms, an aralkyl having 7 to 27 carbon atoms or an acyl having 2 to 22 carbon atoms, x is 0 to 20, m is the valence of K and is an integer from 1 to 6, and y is 1 to 20.

[0018] The magnesium dihalide MgX<sub>2</sub> of said titaniumless complex (a) can be selected from magnesium chloride, magnesium bromide and magnesium iodide. Preferably, it is magnesium dichloride.

[0019] The alkoxy compound K(OR)<sub>m</sub> is in its most general form defined as a component of said titaniumless complex (a), which complex (a) is reacted further with said halogen compound (b) and said titanium halide (c) or said titanium halide (c) and said halogen compound (b). The alkoxy compound is, however, more closely defined in the following description of three main embodiments of the claimed process.

[0020] The gist of the invention is to choose the reactants and their order of reaction so that the magnesium dihalide is present when the titanium halide (c) is reacted.

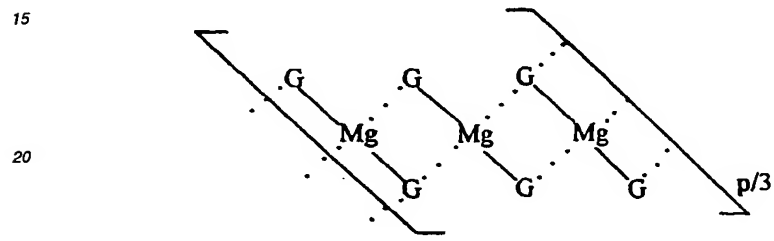
#### First main embodiment

[0021] According to a first main embodiment of the invention, said titaniumless magnesium compound (a) is a compound or complex containing halogen and alkoxide linked to magnesium. It may be a compound such as a magnesium haloalkoxide, typically MgXOR · nR'OH, wherein X is a halogen, preferably chlorine, R and R' are a C<sub>1</sub>-C<sub>12</sub> hydrocarbyl and n is 0-6. Typically, it is a complex product containing magnesium, halogen and alkoxy, essentially characterized by having the following formula (1):

wherein X is a halogen, preferably a chlorine, R is an alkyl group having from 1 to 20 carbon atoms, p is from 2 to 20 and q is < p, preferably < 0.66 p. If there are several halogens X and alkoxy groups OR in the complex product, they can be different or equal.

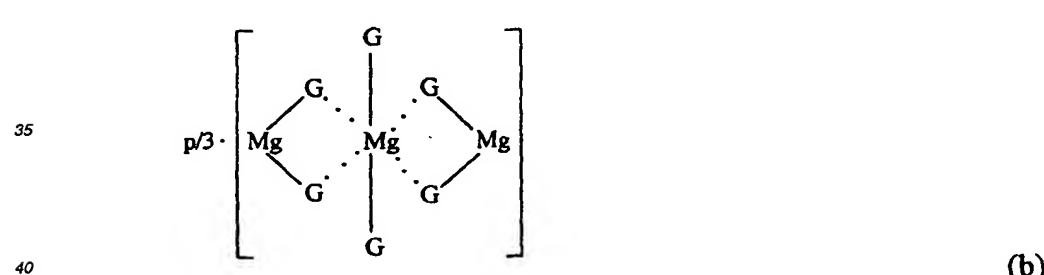
[0022] The complex containing magnesium, halogen and alkoxy used in the invention can, depending on the quality and quantity of elements and groups, preferably be soluble in non-polar organic solvents. The soluble complexes are thus preferably used as starting material for catalytically active stoichiometrical procatalyst complexes. Further, the complex containing magnesium, halogen and alkoxy is always less reductive than magnesium alkyls  $MgR_2$  and  $RMgX$  and is therefore more suitable for activation of the transition metal compound.

[0023] The chemical structure of the titaniumless magnesium compound (a) according to the first main embodiment is based on the bivalence and bridge-forming ability of magnesium. It is believed, without limiting the scope of the invention, that the chemical structure for complexes having  $p \geq 3$  is (a):

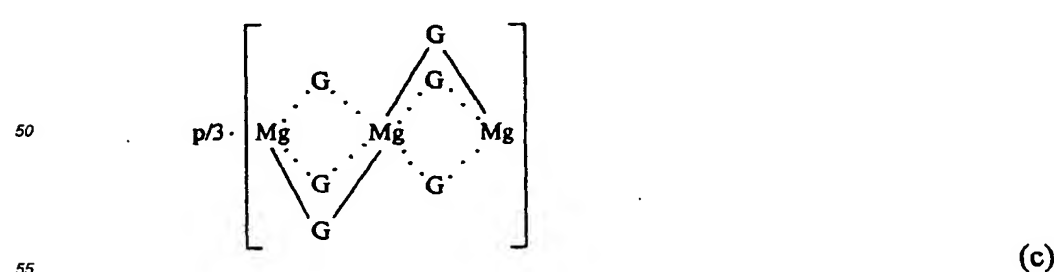


25 wherein each G is the same or different and is selected from said X and said OR to form q units of X and  $2p-q$  units of OR, and p is from 3 to 20. If  $p/3$  is greater than 1 there is in formula (a) a  $\cdots$ -bridge from the furthest Mg—G to the nearest Mg—G of the next unit.

[0024] The chemical structure can also be (b):



45 wherein each G is the same or different and is selected from said X and said OR to form q units of X and  $2p-q$  units of OR, and p is from 3 to 20, or (c):



wherein each G is the same or different and is selected from said X and said OR to form q units of X and  $2p-q$  units of

OR, and p is from 3 to 20.

[0025] When the titaniumless magnesium compound (a) has the composition and structure according to the first main embodiment, the claimed process takes place as follows:

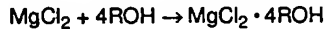
5 (i) said complex containing halogen and alkoxide linked to magnesium as said titaniumless magnesium compound (a) is reacted with said halogen compound (b) to give an intermediate (ab) and then  
 (ii) said intermediate (ab) is reacted with said titanium halide (c), i.e. [(a) + (b)] + (c), or:

10 (i)' said complex containing halogen and alkoxide linked to magnesium as said titaniumless magnesium compound (a) is reacted with said titanium halide (c) to give an intermediate (ac) and then  
 (ii)' said intermediate (ac) is reacted with said halogen compound (b), i.e. [(a) + (c)] + (b).

[0026] In both contact sequences a titaniumless compound containing halogen linked to magnesium is, contrary to Sumitomo, reacted with a titanium halide. When repeating Sumitomo, it gave poorer results than the invention.

15 [0027] In said first main embodiment, said complex is preferably a complex of a magnesium dihalide and a magnesium dialkoxide as said magnesium compound (a). More preferably, it is a magnesium dichloride-magnesium alkoxide complex having the formula  $MgCl_2 \cdot [Mg(OR)_2]_t$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, and t is 1-6, preferably about 2. It is e.g. prepared by reacting magnesium dichloride  $MgCl_2$  with an alcohol ROH into an intermediate which is a magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_t$  and reacting the magnesium dichloride-alcohol complex with t mol of a magnesium dialkyl  $MgR'''_2$ , wherein  $R'''$  is a hydrocarbyl group having 1 to 20 carbon atoms.

20 [0028] Most preferably, the complex of said magnesium dihalide and a magnesium dialkoxide as said alkoxyl compound is a magnesium dichloride-dimagnesium dialkoxide complex having the formula  $MgCl_2 \cdot [Mg(OR)_2]_2$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl. The complex may e.g. be prepared by reacting magnesium dichloride with an alcohol ROH and the obtained intermediate with a dialkyl magnesium  $R'''_2Mg$  essentially as follows:



[0029] In the reaction between the magnesium dihalide, the alcohol and the dialkyl-magnesium, the molar ratio  $MgCl_2$ :ROH is preferably 1:1 to 1:8, most preferably 1:2 to 1:5. The molar ratio  $MgCl_2 \cdot 4ROH$ : $MgR'''_2$  is preferably 1:1 to 1:4, most preferably about 1:2. The temperature is preferably 100 °C to 200 °C and the reaction time preferably about 2 h to about 8 h. A hydrocarbon solvent such as toluene may be present in the reaction.

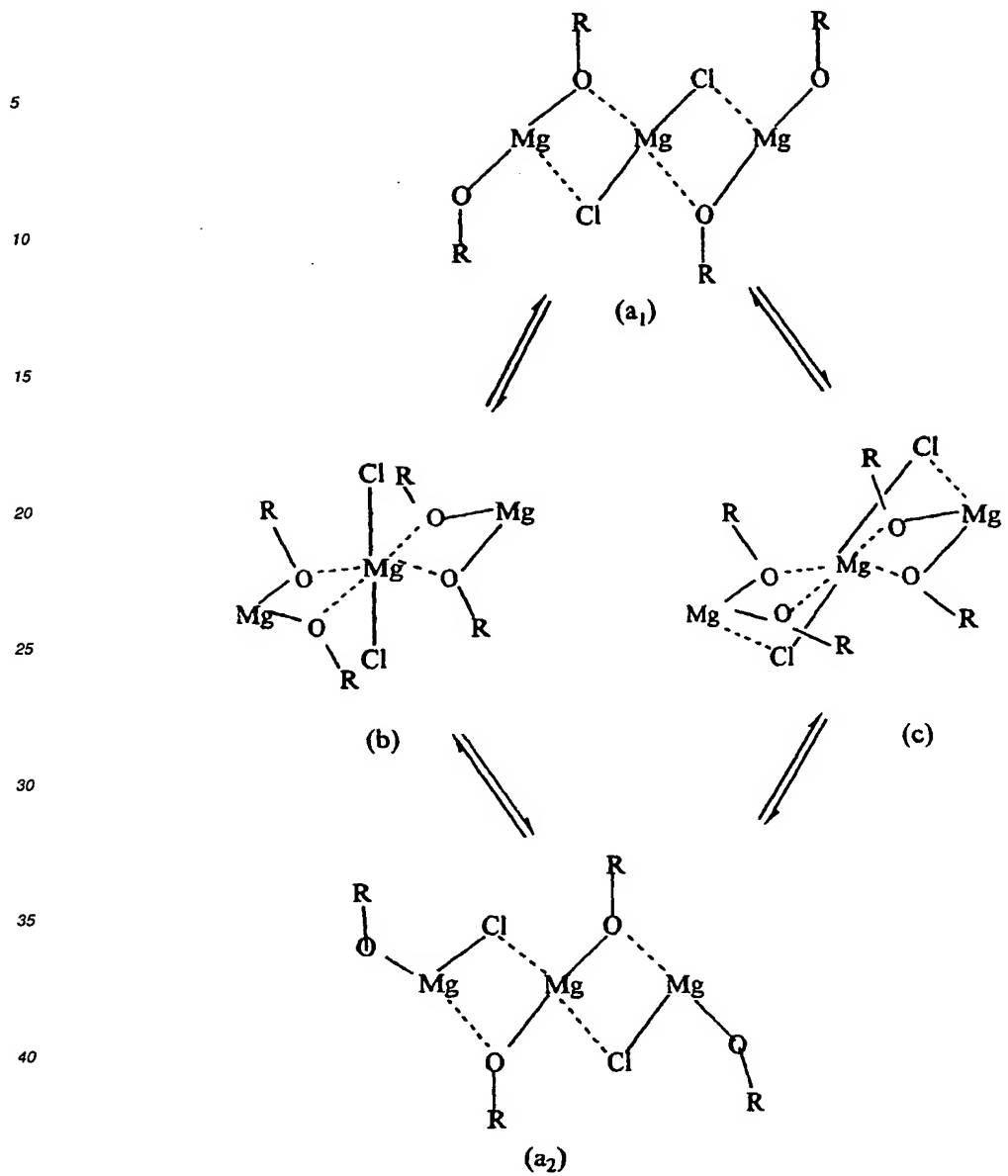
35 [0030] Most probably the complex has the structure of an equilibrium between structures (a<sub>1</sub>), (a<sub>2</sub>), (b) and (c), as illustrated (non-limiting) by the following trimer equilibrium of the  $MgCl_2 \cdot [Mg(OR)_2]_2$  complex:

40

45

50

55



[0031] In the above formulas (a<sub>1</sub>), (a<sub>2</sub>), (b) and (c), Cl can be replaced by any halogen such as fluorine, chlorine, bromine and iodine, but the purposes of the invention are best fulfilled with chlorine.

[0032] The alkyl R of the alkoxy group can be any alkyl suitable for the purpose of the invention. Similar structure and solubility parameter to optional solvents give soluble complexes for stoichiometric preparation of active procatalyst complexes. Different structure and solubility parameters give insoluble complexes for use as activating support. When a solvent having 5-10 carbon atoms, such as toluene, is used, R is preferably an alkyl group having from 1 to 16 carbon atoms, more preferably from 4 to 12 carbon atoms, most preferably from 6 to 10 carbon atoms.

[0033] According to the most preferable variant of said first main embodiment:

(i)' said magnesium dichloride-dimagnesium dialkoxide complex  $MgCl_2 \cdot [Mg(OR)_2]_2$  (a) wherein R is a  $C_1-C_{20}$  alkyl or a  $C_7-C_{27}$  aralkyl, preferably a  $C_6-C_{16}$  alkyl, is reacted with said titanium halide (c), which is said titanium

tetrachloride  $TiCl_4$ , to give an intermediate (ac) and then

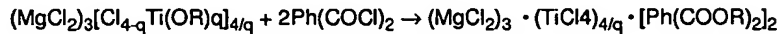
(ii)' said intermediate (ac) is reacted with said halogen compound (b), which is said phthalic acid dichloride  $Ph(COCl)_2$ , wherein Ph is o-phenylene.

5 [0034] The reactions of this variant can, without limiting the scope of protection, e.g. be described by means of the following equations:

Step (1):



Step (2):



15 [0035] Usually, q is 1 or 2. It can e.g. be seen, that the harmful  $Cl_3TiOR$  (when q = 1) after it is formed remains complexed and is finally converted to catalytically active  $TiCl_4$ . No purification or waste problems relating to the formation  $Cl_3TiOR$  arise.

20 [0036] In step (ii) of said first main embodiment, said intermediate (ab) and/or said complex of said magnesium dihalide and a magnesium alkoxide as said magnesium compound (a), respectively, are preferably added to the titanium halide (c), and not vice versa. Even more preferably, they are added drop by drop, to said titanium halide (c). The titanium halide (c) is preferably in liquid form and most preferably hot, such as at 70-140 °C.

25 [0037] In said first main embodiment, the molar ratio between the reactants (a), (b) and (c) is preferably approximately stoichiometric with the exception of the titanium halide (c) in step (ii) which preferably is used in a 5 to 20 fold excess with respect to the magnesium halide. Elevated temperatures are preferably used, whereby said halogen compound (b) preferably is reacted at 50 °C to 75 °C and said titanium halide (c) in step (ii) preferably is reacted at 70 °C to 110 °C.

30 [0038] Although the first main embodiment reduces the need for repeated titanations and washes, a still more active and pure product is obtained if the titanation and hydrocarbon wash are repeated 1 to 3 times.

30 [0039] An example of the first main embodiment is presented in an enclosed scheme, see Figure 1.

### Second main embodiment

35 [0040] According to a second main embodiment of the invention, said titaniumless magnesium compound (a) is a complex of said magnesium dihalide and an alcohol carrying said alkoxy moiety, or, alternatively, a non-complex magnesium dialkoxide.

[0041] The claimed process then takes place as follows:

40 (i) said titaniumless magnesium compound (a) which is selected from said complex of said magnesium dihalide and said alcohol, and said non-complex magnesium dialkoxide, is reacted with said halogen compound (b) to give an intermediate (ab) which is a complex of a magnesium dihalide and said electron donor and  
(ii) said intermediate (ab) which is a complex of said magnesium dihalide and said electron donor is reacted with said titanium halide (c).

45 [0042] In a first variant of said second main embodiment, a non-complex magnesium dialkoxide is used as the starting material (a) of step (i). It is preferably a magnesium dialkoxide  $Mg(OR)_2$ , wherein R is a  $C_1-C_{20}$  alkyl or a  $C_7-C_{27}$  aralkyl, preferably a  $C_6-C_{16}$  alkyl. It can be prepared by any suitable process such as the process described by Sumitomo, see column 9, line 56, to column 10, line 13. In the present invention, however, it is preferably prepared by reacting a magnesium dialkyl and an alcohol ROH. The reaction can e.g. be described by means of the following equation:



wherein R and R''' are as defined above.

55 [0043] In the first variant of said second main embodiment the most preferable process comprises the steps wherein:

(i) said titaniumless magnesium compound (a), which is said magnesium dialkoxide  $Mg(OR)_2$ , wherein R is a  $C_1-C_{20}$  alkyl or a  $C_7-C_{27}$  aralkyl, preferably a  $C_6-C_{16}$  alkyl, is reacted with said halogen compound (b), which is said phthalic acid dichloride  $Ph(COCl)_2$ , wherein Ph is o-phenylene, to give an intermediate (ab) which is a complex of

EP 0 926 165 A1

said magnesium dichloride and said phthalic acid diester  $\text{Ph}(\text{COOR})_2$ , and

(ii) said intermediate (ab), which is said complex of said magnesium dichloride and said phthalic acid diester  $\text{Ph}(\text{COOR})_2$ , is reacted with said titanium halide (c), which is said titanium tetrachloride  $\text{TiCl}_4$ .

5 [0044] The reactions can, without limiting the scope of protection, e.g. be described by means of the following equations:

Step (i):



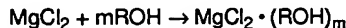
and

Step (ii):



wherein  $m'$  is about 1 to about 2.

20 [0045] In a second variant of said second main embodiment, a complex of said magnesium dihalide and an alcohol is used as said titaniumless magnesium compound (a) of step (i). It is preferably a magnesium dichloride alcohol complex  $\text{MgCl}_2 \cdot (\text{ROH})_m$ , wherein R is a  $\text{C}_1\text{-C}_{20}$  alkyl or a  $\text{C}_7\text{-C}_{27}$  aralkyl, preferably a  $\text{C}_6\text{-C}_{16}$  alkyl, and m is 1-6. The complex is preferably prepared by reacting magnesium dichloride and an alcohol, e.g. as illustrated by the following equation:



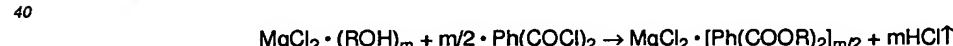
25 [0046] In the second variant of said second main embodiment, the most preferable process then comprises the steps wherein:

30 (i) said titaniumless magnesium compound (a), which is said magnesium dichloride-alcohol complex  $\text{MgCl}_2 \cdot (\text{ROH})_m$ , wherein R is a  $\text{C}_1\text{-C}_{20}$  alkyl or a  $\text{C}_7\text{-C}_{27}$  aralkyl, preferably a  $\text{C}_6\text{-C}_{16}$  alkyl, and m is 1-6, is reacted with said halogen compound (b), which is said phthalic acid dichloride  $\text{Ph}(\text{COCl})_2$ , wherein Ph is o-phenylene, to give a intermediate (ab), which is a complex of said magnesium dichloride and said phthalic acid diester  $\text{Ph}(\text{COOR})_2$ , wherein R is the same as above, and

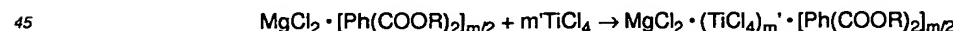
35 (ii) said intermediate (ab) which is said complex of said magnesium dichloride and said phthalic acid diester  $\text{Ph}(\text{COOR})_2$ , is reacted with said titanium halide (c), which is said titanium tetrachloride  $\text{TiCl}_4$ .

[0047] The reactions can, without limiting the scope of protection, be described by means of the following equations:

Step (i):



Step (ii):



wherein m is 1-6 and  $m'$  is from 0.5 to m.

50 [0048] In step (ii) of said second main embodiment, said complex of said magnesium dihalide and said electron donor is preferably added to the titanium halide (c), and not vice versa. Even more preferably, it is added drop by drop, to said titanium halide (c). The titanium halide (c) is preferably in liquid form and most preferably hot, such as at 70-140 °C.

[0049] In said second main embodiment, the molar ratio between the reactants (a), (b) and (c) is approximatively stoichiometric, preferably with the exception of the titanium halide (c) which more preferably is used in a 5 to 20 fold excess with respect to the magnesium halide. Elevated temperatures are preferably used, whereby said halogen compound (b) preferably is reacted at 50 °C to 75 °C and said titanium halide (c) preferably is reacted at 70 °C to 110 °C.

55 [0050] Although said second embodiment reduces the need for repeated titanations and washes, a still more active and pure product is obtained if the titration and hydrocarbon wash are repeated 1 to 3 times.

[0051] An example of the second main embodiment is presented in an enclosed scheme, see Figure 2.

## Third main embodiment

[0052] According to a third main embodiment of the invention, said titaniumless magnesium compound (a) is a complex of said magnesium dihalide and an alcohol carrying said alkoxy group.

5 [0053] In that case:

(i)' said titaniumless magnesium compound (a) which is said complex of said magnesium dihalide and an alcohol, is reacted with said titanium halide (c), to give an intermediate (ac), and  
 (ii)' said intermediate (ac) is reacted with said halogen compound (b).

10 [0054] In the third main embodiment, the complex of said magnesium dihalide and an alcohol is preferably a magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_m$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl and m is 1-6. It can be prepared as described above.

[0055] According to the most preferable variant of said third embodiment, the process comprises the following steps:

15 (i)' titaniumless magnesium compound (a), which is said magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_m$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl and m is 1-6, is reacted with said titanium dihalide (c), which is said titanium tetrachloride  $TiCl_4$ , to give an intermediate (ac) and  
 20 (ii)' said intermediate (ac) is reacted with said halogen compound (b), which is said phthalic acid dichloride  $Ph(COCl)_2$ , wherein Ph is o-phenylene.

[0056] This reaction can, without limiting the scope of protection, e.g. be described by means of the following equation:

25 Step (i)':



Step (ii)':



wherein m is from about 1 to about 6, preferably about 2, q is 1-4, preferably about 2, and R is as said above.

[0057] In step (i)' of said third main embodiment, said titanium halide (c) is preferably added to said complex of said magnesium dihalide and an alcohol as said titaniumless magnesium compound (a), and not vice versa. Even more preferably, it is added drop by drop, to said titaniumless magnesium compound (a). The titanium halide (c) is preferably in liquid form.

[0058] In said third main embodiment, the molar ratio between the reactants (a), (b) and (c) is preferably approximately stoichiometric.

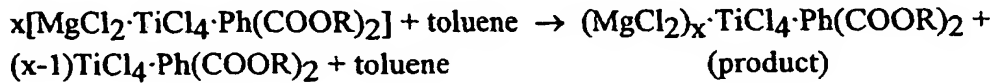
40 [0059] Although said third embodiment reduces the need for repeated titanation and washes, a still more active and pure product is preferably obtained if the final titanation and hydrocarbon (e.g. toluene) wash are repeated 1 to 3 times.

[0060] An example of the third main embodiment is presented in an enclosed scheme, see Figure 3.

[0061] As was stated above, common to the first and second main embodiments is, that in step (ii) said intermediate (ab) is preferably added, more preferably drop by drop, to said titanium halide (c), which is in liquid form and preferably hot, more preferably at 75-150 °C. Further, all embodiments may further comprise a step wherein:

(iii) the obtained product of step (ii) or (ii)' is further treated with said titanium halide (c) and/or washed, preferably repeatedly washed with an aromatic hydrocarbon, such as toluene, or an organic liquid having the same solubility parameter as said aromatic hydrocarbon.

50 [0062] The washing can, without limiting the scope of protection, be illustrated by the following equation:



x is a number larger than 1.

[0063] It is seen that the MgCl<sub>2</sub> is concentrated in the catalyst component, which leads to higher activity.

#### Product and use

[0064] In addition to the above described process, the invention also relates to a catalyst component comprising magnesium, titanium, a halogen and an electron donor. The catalyst component is characterized in that it has been prepared by the process described in any of claims 1 to 15 or in the preceding text. Preferably, the claimed catalyst component is an isolated complex of a magnesium dihalide, preferably magnesium dichloride, an electron donor obtained by replacing the halogen of a halogen compound by an alkoxy group, preferably a phthalic acid diester, and a titanium halide, preferably titanium tetrachloride. Preferably, said complex is prepared by contacting stoichiometric amounts of said components (a), (b) and (c). Preferably, said complex has an X-ray pattern comprising a peak between 5° and 10° 2θ (Siemens D500 instrument, CuK $\alpha$  radiation wavelength 1.541 Å, effect 40 kV and 35 mA). Most preferably, said complex has an X-ray pattern comprising a crystal height indicating peak between 16° and 18° 2θ.

[0065] The invention also relates to the use of said catalyst component for the polymerization of  $\alpha$ -olefins, preferably propene. In such a polymerization, said catalyst component is preferably used together with another catalyst component comprising an organometal compound of a metal belonging to Group 1, 2 or 13 (IUPAC 1990) of the Periodic Table, preferably an alkyl aluminium compound, is used. The organometal compound is in the art called a cocatalyst. Further, another electron donor may be used together with said catalyst component and the co-catalyst. Such a donor is in the art called an external electron donor.

#### Examples 1 to 4 (first main embodiment)

##### Preparation of the catalyst component complex

[0066] 1.69 g (17.70 mmol) of anhydrous MgCl<sub>2</sub> was introduced in inert conditions into a 100 ml septum bottle. 11.12 ml (9.27 g, 70.80 mmol) of 2-ethyl-hexanol (EHA) was introduced on to the MgCl<sub>2</sub> and after this the temperature was increased to 125-128 °C to allow the reaction components to react with each other. After this, 8.81 ml (7.67 g, 83.19 mmol) of toluene was added after the reaction solution had cooled down to 110 °C. After the addition of the toluene the reaction solution was cooled down to 21 °C. Then 40 ml (29.16 g, 35.4 mmol) of a 20 w-% heptane solution of butyl-octyl-magnesium (BOMAG) was added. After this 5.10 ml (7.19 g, 35.4 mmol) of phthaloyl dichlorid (PDC) was added to produce a MgCl<sub>2</sub> donor complex solution.

[0067] The MgCl<sub>2</sub> donor complex was now, drop by drop, added into 38.91 ml (67.16 g, 354 mmol) of TiCl<sub>4</sub> and allowed to react with this reagent at a temperature of 95 °C. The reactants were allowed to react with each other for 30 min.

[0068] After the TiCl<sub>4</sub> treatment, the complex was allowed to settle and the liquid was siphoned off. After this, 100 ml (86.6 g, 0.94 mol) of toluene was added on to the complex and the complex was washed in this solution at 90 °C for 20 min. Depending on which of the synthesis was under work, this washing step was done once (example 1), twice (example 2), three times (example 3) or four times (example 4). Finally, the catalyst complex was washed twice with 65 ml (44.44 g, 0.44 mol) portions of heptane for 20 min at 80 °C and thereafter, the complex was washed at room temperature with a 55 ml (34.44 g, 0.48 mol) portion of pentane for 20 mm to improve the drying conditions. The catalysts were dried under a stream of nitrogen for one hour.

##### Chemical characterization of the complexes

[0069] The catalyst complexes were characterized with respect to their chemical composition by measuring their Ti and Cl content. The Ti analysis was started by dissolving the samples in a mixture of nitric and hydrofluoric acid. The metal was measured flame atomic absorption with a nitrous acetylene flame. Chloride was determined after dissolution in dilute sulphuric acid by potentiometric titration with a standard silver nitrate solution.

##### Determination of donors and phthalic anhydride

[0070] The determination of the phthalic esters and the phthalic anhydride were done by first dissolving the sample in acetone. The dissolving was improved by keeping the acetone slurry in an ultra-sound bath for 5 min. After this the samples were filtered and run by solution chromatography. As eluent a solution consisting of water and acetonitril in the proportion of 4/96 was used. Eluent flow rate was 1.5 ml/min. A photo diode array was used as detector. Each component was identified by comparing the respective retention time and UV spectra with standard components.

## GC studies to measure alcohol content

[0071] To check the conversion rate of the ethanol (EtOH), 2-ethyl-hexanol (EHA), or other alcohol added in the synthesis, the alcohol content of the catalysts were measured by gas chromatography (GC). This was done by first dissolving a 100 mg sample of the catalyst in 1 ml of n-pentanol. Depending on the alcohol to be measured, an internal alcohol standard was chosen. If ethanol was to be measured the n-pentanol solution contained n-propenol as internal standard. To improve the solubility of the catalyst in the solution, the sample was kept in an ultra-sound bath. To remove the inorganics from the organic solution it was extracted with 1 ml of water and to ensure full dissolution, another ml of the n-pentanol solution was added. To ensure repeatable equilibrium conditions between the organic layer and the water layer the samples were allowed to stand overnight. The sample for the GC was taken from the alcohol layer. A Hewlett Packard 5890 GC with a 60 m DB-1 column was used for the GC analyses. The column had a diameter of 0.25 mm with a film thickness of 1  $\mu$ m. An FID detector was used.

## Bulk polymerization

[0072] Propylene was polymerized in stirred tank reactor having a volume of 5 l. About 0.9 ml triethyl aluminium (TEA) as a cocatalyst, ca 0.12 ml of a 100-% solution of cyclohexyl methyl dimethoxy silane as an external donor and 30 ml of n-pentane were mixed and allowed to react for 5 minutes. Half of the mixture was added to the polymerization reactor and the other half was mixed with ca 20 mg of a catalyst complex. After additional 5 minutes the catalyst/TEA/donor/n-heptane mixture was introduced into the reactor. The Al/Ti mole ratio was 250 and the Al/external donor mol ratio was 10 mol/mol. 70 mmol hydrogen and 1400 g of propylene were introduced into the reactor and the temperature was raised within 15-30 minutes to 70 °C. The polymerization time was 60 minutes, after which the polymer formed was taken out from the reactor. The polymers were characterized with respect to their Melt Flow Rate (MFR<sub>2</sub>), bulk density (BD) and fraction of total solubles in xylene (TS).

## Results

## Preparation of the complexes

[0073] The catalyst complexes achieved in this investigation are listed in Table 1.

Table 1

The catalyst complexes prepared.			
Example	Number of toluene washes	Colour of catalyst	Morphology of catalyst
1	1	Dark wine-red	Freely flowing
2	2	Dark wine-red	Freely flowing
3	3	Dark wine-red	Freely flowing
4	4	Dark wine-red	Freely flowing

## The chemical composition of the catalysts

[0074] The chemical composition of the catalysts were measured according to the description in the experimental section. In Table 2 the chemical composition of the catalysts are listed in w-% units, in Table 3 the composition is listed in mol-% units and in Table 4 the molar proportions between Mg, Ti and DOP are compared.

[0075] With three washes a composition of  $(\text{MgCl}_2)_6\text{TiCl}_4\text{DOP}$  was achieved. During the washes, there was a slightly higher wash out of  $\text{TiCl}_4$  compared to DOP in the last catalyst. The amount of free alcohol (EHA) was also very low playing no significant part in the chemical composition (now 0.004-0.006 mol-%), i.e. being about 5% of the mol amount of  $\text{TiCl}_4$  or DOP. The amount of phthalic anhydride was about 50% of the DOP amount. To sum up the results from the chemical measurements it can be said that the chemical composition of the catalyst complex when using the  $\text{MgCl}_2$  enriched  $\text{Mg}(\text{OR})_2$  as a reagent in the catalyst synthesis is  $(\text{MgCl}_2)_3\text{TiCl}_4\text{DOP}(\text{PA})_{0.5}$ .

EP 0 926 165 A1

Table 2

The chemical composition of the catalysts in w-% units					
Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
1	7.8	4.7	33.6	0.72	6.7
2	8.1	4.7	32.0	0.54	7.4
3	10.2	3.2	28.5	0.58	6.5
4	12.9	1.6	21.6	0.51	6.1

Table 3

The chemical composition of the catalysts in mol-% units					
Example	Mg mol-%	Ti mol-%	DOP mol-%	EHA mol-%	PA mol-%
1	0.321	0.098	0.086	0.0055	0.045
2	0.333	0.098	0.082	0.0041	0.050
3	0.420	0.067	0.073	0.0045	0.044
4	0.531	0.033	0.056	0.0039	0.041

Table 4

The molar composition between Mg, Ti and DOP			
Example	Mg	Ti	DOP
1	3.3	1	0.9
2	3.4	1	0.8
3	6.3	1	1.1
4	15.9	1	1.7

**Calculated and found chlorine contents**

[0076] The chlorine content in the catalysts were calculated on the basis of the Mg and Ti content. The calculations were based on the assumption the Mg was present in the catalyst as  $MgCl_2$  and Ti as  $TiCl_4$ . These calculated results were then compared to the measured results. The results are listed in Table 5. The results showed to be in good agreement, which indicates that both Mg and Ti are present in the catalyst complexes in the fully chlorinated form.

Table 5

The calculated and the found chlorine content in the catalysts		
Example	Calculated Cl w-%	Found Cl w-%
1	36.7	36.9
2	37.6	38.0

Table 5 (continued)

The calculated and the found chlorine content in the catalysts		
Example	Calculated Cl w-%	Found Cl w-%
3	39.3	39.7
4	42.4	43.8

10 Wash out of  $\text{TiCl}_4 \cdot \text{DOP}$ 

[0077] All the chemical measurements support the same conclusion: due to the toluene  $\text{TiCl}_4$  and DOP are washed out from the catalyst in a molar proportion of 1:1. This shows up as a constant decrease of the Ti mol-% and the DOP mol-%, and as a constant increase of the Mg mol-% and the Cl mol-%.

## 15 Activity of the catalysts

[0078] All the catalyst complexes were test polymerized according to the descriptions in the experimental section. The results are listed in Table 5. The results showed that all the catalyst complexes had about the same activity, being 20 between 1.0 and 1.5 kg PP/g cat.

Table 6

The test polymerization results		
Example	Activity kg PP/g cat	Activity kg PP/g Ti
1	1.1	23
2	1.2	26
3	1.5	45
4	1.3	81

## 35 MFR of the polymers

[0079] In Table 7 the MFR values achieved from the test polymerization results are listed. The results indicated a systematic increase in MFR with increasing number of toluene washes as MFR increases from 2.0 in the first polymer to 13.7 in the third.

## 40 Table 7

The MFR values of the polymers	
Example	MFR
1	2.0
2	4.9
3	13.7
4	12.4

## 50 Examples 5 to 9 (second main embodiment)

## 55 Preparation of the catalyst component complexes

[0080] All chemicals were handled in strict inert conditions and all the reactions took place also in strict inert conditions in nitrogen atmosphere.

[0081] 8.85 mmol of butyl-octyl-magnesium was introduced into a 150 ml glass reactor. A 20% heptane solution (BOMAG-A) was used giving a feed volume of 10 ml (7.29 g). 17.7 mmol (2.78 ml, 2.32 g) of 2-ethyl-1-hexanol (EHA) was then added at room temperature. The temperature was increased to 60 °C and the reactants were allowed to react with each other at that temperature for 30 min. After this 8.85 mmol (1.28 ml, 1.80 g) of phthaloyl chloride (PDC) was added and the reactants were again allowed to react with each other for 30 min at 60 °C. The resulting solution was added dropwise into 88.5 mmol (9.73 ml, 16.79 g) of  $TiCl_4$  that had been preheated to 95 °C. The reactants were also in this case allowed to react with each other for 30 min at 95 °C. After this 60 ml of toluene was added. After the precipitate had settled the mother liquid was siphoned off. Five different examples were carried out according to this description. After this the catalyst complex was washed with 30 ml portions of toluene. In example 5, the complex was washed once with toluene, in example 6 twice, in example 7 three times, in example 8 four times and in example 9 six times with 30 ml portions of toluene. The toluene washes were carried out at 90 °C. Last the complex was washed three times with 30 ml portion of pentane. The complexes were finally dried under a stream of nitrogen. The yield of the catalyst was about 2 g which corresponded to about 75% of the theoretical.

15 **Characterization of the catalyst components**

[0082] The catalyst component complexes were analyzed with respect to their Mg, Cl and Ti content. In addition to this, the amount of donor compound, the di-octyl-phthalate (DOP) formed in the synthesis, was measured from the catalysts. To indicate to what degree the formed donor compound (DOP) was decomposing in the synthesis, the amount 20 of phthalic anhydride (PA) was also measured from the catalysts.

IR and X-ray of the unwashed Mg:Ti:DOP complex

[0083] A stoichiometric complex of  $MgCl_2 \cdot TiCl_4 \cdot DOP$  was prepared by reacting 6.37 mmol (7.19 ml, 5.24 g) of BOMAG with 12.729 mmol (2.00 ml, 1.67 g) of EHA in a 50 ml glass reactor. After this 6.37 mmol (0.92 ml, 1.29 g) of phthaloyl chloride was introduced and last 6.37 mmol (0.70 ml, 1.21 g) of  $TiCl_4$  was added. The solid product was washed with pentane and finally, the sample was dried in a stream of nitrogen. The sample was characterized by IR spectroscopy and by means of its X-ray diffraction pattern.

30 **The IR studies**

[0084] IR spectra were taken by means of a Nicolet 510 FTIR equipment with  $2\text{ cm}^{-1}$  resolution. The number of scans were 128. All the samples were investigated as capillary films between two KBr tablets. The pure EHA was not handled in inert conditions, while the  $MgCl_2$  samples were handled in a glove box in an inert nitrogen environment in order to 35 protect the samples from air and moisture.

X-ray diffraction patterns

[0085] The WAXS patterns were collected in a reflection mode between 2° and 70° 2θ with a Siemens D500 instrument. The diffractometer was equipped with a Cu anode and a graphite monochromator in the reflected beam. The CuK $\alpha$  radiation wave-length was 1.541 Å. The effect used was 40 kV and 35 mA. The sample was loaded in a glovebox into a Mylar film covered sample holder.

45 **Bulk polymerization**

[0086] The bulk test polymerization was carried out according to the description on page 18.

**Results**

50 **Preparation of the complexes**

[0087] The reaction between the Mg-alkyl and the alcohol resulted in a clear solution with a little bit higher viscosity. The reaction was exothermic as the solution became warm when mixing the reactants, the temperature increase was from room temperature up to 50 °C. When the phthaloyl chloride was added a slight yellow colour appeared. Also this 55 reaction was slightly exothermic. The reaction solution became again freely flowing with a low viscosity.

[0088] The  $TiCl_4$  was introduced into a 150 ml glass reactor and heated to 95 °C. The Mg solution was then added to the hot  $TiCl_4$  solution dropwise. A beige precipitate started to form right at the beginning of the addition. During addition the solution turned turbid. A partly freely floating precipitate was formed together with more tarlike precipitate that

EP 0 926 165 A1

started to foul the reactor walls. To improve the settling conditions toluene was added to the reaction solution. A satisfactory settling of the product was then achieved so that the reaction solution could be siphoned off. Depending on the number of toluene washes the resulting product became more freely flowing. If only one toluene wash was used the product was still as agglomerates, but already two toluene washes resulted in a freely flowing powder-like product.

5 [0089] In the case of the catalyst components of examples 5, 6 and 7, a joined  $MgCl_2$ -DOP complex and a joined addition to the  $TiCl_4$  solution was carried out. After the first toluene wash, 1/3 of the solution slurry was separated. The separated part was then washed with the aliphatic hydrocarbon and dried to give the product of example 5. The remaining part of the slurry was washed a second time with toluene and half of this solution slurry was then taken out from the reactor and undertaken the same hydrocarbon treatment as in example 5, resulting in the product of example 6. The 10 remaining part of the catalyst slurry in the reactor was washed twice with toluene and then washed with an aliphatic hydrocarbon in the same way as the first two examples. This sample was the product of example 7. The catalyst morphologies are listed in Table 8.

15

Table 8

The morphology of the catalysts		
Example	Number of toluene washes	Morphology of catalyst
5	1	Black agglomerates
6	2	Dark powder
7	4	Dark powder

25

**The chemical composition of the catalysts**

[0090] The Mg, Ti, Cl, DOP, EHA and the phthalic anhydride (PA) content of the catalysts were measured. The results are listed in w-% units in Table 9. In Table 10 the chemical composition is given in mol-% units and in Table 11 the Mg and DOP amounts are compared to the Ti amount on a molar basis. Table 12 shows the Cl content of the catalysts.

35

Table 9

The chemical composition of the catalysts in w-% units					
Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
5	5.7	6.8	47.7	0.26	3.4
6	11.3	3.1	32.2	0.18	2.5
7	13.4	1.4	21.3	0.25	1.7

40

45

Table 10

The chemical composition of the catalysts in mol-% units					
Example	Mg mol-%	Ti mol-%	DOP mol-%	EHA mol-%	PA mol-%
5	0.235	0.142	0.122	0.002	0.023
6	0.465	0.065	0.083	0.001	0.017
7	0.551	0.029	0.055	0.002	0.012

55

Table 11

The molar proportions between Mg and Ti and between DOP and Ti			
Example	Mg	Ti	DOP
5	1.7	1	0.86
6	7.2	1	1.28
7	18.9	1	1.87

Table 12

The calculated Cl content in the catalysts compared to the measured amounts		
Example	Calculated Cl w-%	Found Cl w-%
5	36.8	36.4
6	42.2	45.0
7	43.3	44.5

#### Activity of the catalysts

[0091] All the catalysts were test polymerized according to the above instructions. The polymerization results are listed in Table 13 in both kg PP/g cat ad kg PP/g Ti units. The activities are also shown in Figure 4 and Figure 5. Activities of almost 8 kg PP/g cat were reached. Catalysis of the examples 5, 6 and 7 gave good polymerization results, with the highest activity achieved for the catalyst that had been twice washed with toluene. The activities expressed in kg PP/g Ti units showed a linear increase related to the number of toluene washes for the catalysts of examples 5, 6 and 7 (Figure 5). Activities of over 500 kg PP/g Ti were reached.

#### Characterization of the polymers

[0092] All the polymers were characterized with respect to their melt flow rate (MFR) and bulk density (BD). All the polymers showed to have a MFR<sub>2</sub> between 11-12 g/10 min, indicating a quite good hydrogen response. Bulk densities were between 0.35-0.39 g/ml. The total solubles were between 2 and 3%, being better for the polymers achieved with the catalyst giving higher activity. The results listed in Table 14.

Table 13

The polymerization results		
Example	Activity kg PP/g cat	Activity kg PP/g Ti
5	2.6	38
6	7.9	254
7	7.3	524

Table 14

The polymer properties			
Example	MFR 2.16 kg, 10 min	TS %	BD g/ml
5	11.0	3.1	0.360
6	12.4	2.1	0.350
7	11.0	2.1	0.390

#### IR studies of the catalyst

15 [0093] IR spectra in the corresponding regions of 1500-1950  $\text{cm}^{-1}$  (Figure 6) and of 1000-1450  $\text{cm}^{-1}$  (Figure 7) were taken from the resulting catalyst of example 6 and compared to an IR spectrum of a typical active catalyst complex coming from a synthesis starting from a  $\text{MgCl}_2 \cdot (\text{EtOH})_3$  support material. The spectra are essentially different, and also different from the IR spectra of the isolated complexes of  $\text{TiCl}_4/\text{DOP}$  and  $\text{MgCl}_2/\text{DOP}$ .

#### X-ray studies of the catalysts

20 [0094] As described above, X-ray diffraction patterns were taken from the resulting catalysts and compared to a X-ray pattern from an inactive catalyst complex and a typical active catalyst complex prepared from a  $\text{MgCl}_2 \cdot 3\text{EtOH}$  support material.

25 [0095] In Figure 8 are shown the X-ray patterns of  $\text{Mg}(\text{OR})_2$  (A)  $\text{MgCl}_2 \cdot \text{TiCl}_4 \cdot \text{DOP}$  (B) produced from  $\text{MgCl}_2 \cdot 3\text{EtOH}$ , and of  $(\text{MgCl}_2)_{1.7} \cdot \text{TiCl}_4 \cdot \text{DOP}$  (C) produced by adding one mol  $\text{MgCl}_2 \cdot \text{DOP}$  to 10 moles of  $\text{TiCl}_4$ . In all these cases there was a strong peak located somewhere between  $5^\circ$  and  $9^\circ 2\theta$ . In addition, there seems to be a halo formation between  $17^\circ$  and  $23^\circ 2\theta$ . The strong peak in the left corner of the pattern indicates that large organic groups are separating metal layers at a distance of between 9 and 17  $\text{\AA}$ , the distance depending on the size of the 30 organic compound (DOP or di-undecyl phthalate DUP). It can thus be stated that the X-ray diffraction patterns for the catalyst complexes originating from the claimed process all show unique features originating from the starting compounds of  $\text{Mg}(\text{OR})_2$  and  $\text{MgCl}_2 \cdot \text{DOP}$ . These patterns show almost no sign of amorphous or crystalline  $\text{MgCl}_2$ .

#### Stoichiometric comparison

35 [0096] To get a fair picture about how much more stoichiometric the claimed process is compared to a conventional Ziegler-Natta PP catalyst component process, the waste amounts and the volumes of chemicals to be circulated are listed in Table 15. As reference is used a classical Z-N PP catalyst component synthesis with two titanations followed by three heptane washes (as in EP 0 491 566). Example 6 was chosen as the best representative. Here two toluene 40 washes has been used to purify the catalyst. As can be seen from the list, the most essential difference between these two synthesis routes is the lack of the titanium alkoxy trichloride waste material, the  $\text{ORTiCl}_3$ , in this new recipe. The lack of Ti waste material makes a great difference in easiness in circulating  $\text{TiCl}_4$ . The other significant change is the decrease in the overall use of  $\text{TiCl}_4$ , that has dropped to one fourth of what it has been in the classical recipe. The aliphatic hydrocarbon wash in the classical recipe have been changed to a toluene dito in the new recipe.

Table 15

Stoichiometric comparison between a classical Ziegler-Natta PP catalyst synthesis and the claimed (example 6) catalyst synthesis. The figures refer to mol/mol Mg.		
Species	Classical Z-N PP cat synthesis	Example 6
$\text{Cl}_3\text{TiOR}$ waste to be neutralized	3	0
$\text{TiCl}_4$ to be circulated	40	10
Donor in excess	0.1	0.8
Hydrocarbon to be circulated	40	7

Table 15 (continued)

5 Stoichiometric comparison between a classical Ziegler-Natta PP catalyst synthesis and the claimed (example 6) catalyst synthesis. The figures refer to mol/mol Mg.

Species	Classical Z-N PP cat synthesis	Example 6
Toluene to be circulated	0	60

## 10 Examples 8 (second main embodiment), 9 and 10 (third main embodiment)

15 [0097] The following reagents are used;  $MgCl_2$  or  $MgR_2$ , 2-ethyl-hexanol (EHA), phthaloyl dichloride (PDC) and  $TiCl_4$  and they are added in the molecular proportion of 1:2:1:1. In the first synthesis (example 8), the Mg-alkyl is reacted with the alcohol, then the phthaloyl chloride (PDC) is added and finally the  $TiCl_4$  is added. In the next two syntheses (examples 9 and 10), the Mg-alkyl is replaced by  $MgCl_2$ . Either the  $TiCl_4$  or the phthaloyl chloride is added in the next step, followed by the last reagent. The synthesis set-ups are listed in Table 16.

Table 16

20 Addition order of the reaction components in the catalyst synthesis

Reaction component/Example	8	9	10
$MgR_2$	1		
$MgCl_2$		1	1
R'OH	2	2	2
PDC	3	3	4
$TiCl_4$	4	4	3

## 35 Preparation of the complexes

40 [0098] The same volumes of reagents have been used in all the experiments regardless in which order they have been added. Thus 22.22 mmol (25.10 ml, 18.3 g) of a 20% heptane solution of butyl-octyl-Mg (BOMAG) was added in experiment (8) and 22.60 mmol (2.15 g) of  $MgCl_2$  was added in experiment (9) and (10). To this, 45.19 mmol (7.10 ml, 5.92 g) of 2-ethyl-1-hexanol EHA was added. The  $TiCl_4$  mol amount added was equal to the mol amount of  $MgCl_2$  being 22.60 mmol (2.48 ml, 4.29 g) and also equal to the mol amount of PDC added, which was 22.60 mmol (3.26 ml, 4.59 g). The addition orders of the reaction components in each catalyst synthesis are listed in Table 16. All the complexes were washed three times with a 100 ml portion of heptane at 90 °C for 15 min and last with a 100 ml portion of pentane at room temperature. Finally the catalysts were dried under a stream of nitrogen.

## 45 Characterization of the catalysts

50 [0099] All the catalysts were characterized with respect to their chemical composition by measuring their Mg, Ti, Cl and di-octyl-phthalate (DOP) content. The Ti and Mg containing catalyst samples were dissolved in a mixture of nitric and hydrofluoric acid and the metals were measured by flame atomic absorption with a nitrous oxide/acetylene flame. Chloride was determined after dissolution in dilute sulphuric acid by potentiometric titration with a standard silver nitrate solution.

55 [0100] The determination of the phthalic esters and the phthalic anhydride were done by first dissolving the sample in acetone. The dissolution was improved by keeping the acetone slurry in an ultra-sound bath for 5 min. After this the samples were filtered and run by solution chromatography. As eluent a solution consisting of water and acetonitrile in a proportion of 4/96 was used. The eluent flow rate was 1.5 ml/min. A photo diode array was used as detector. Each component was identified by comparing its retention time and UV spectra with those of standard components. To further characterize the complexes, IR spectra and X-ray diffraction patterns were taken of them.

EP 0 926 165 A1

**Bulk polymerization**

[0101] The polymerization was carried out as before. See page 18.

5 **Results**

**Chemical composition of the catalysts**

[0102] As stated in the experimental section, the catalysts were characterized with respect to their chemical composition. In Table 17 the chemical composition of the catalysts with respect to the Mg, Ti, di(2-ethyl-1-hexyl)phthalate (DOP), 2-ethyl-1-hexyl alcohol (EHA) and phthalic anhydride PA contents are listed in w-% units and in Table 18 the same species are listed in mol-% units and last; in Table 19 the molar composition between Mg, Ti and DOP are listed. The examples 8 and 10 are represented by two catalysts, 8a and 8b, as well as 10a and 10b, respectively. The chlorine contents are listed in Table 20.

15

Table 17

The Mg, Ti, DOP, EHA and PA contents of the catalysts in w-% units					
Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
8a	3.6	5.6	35.6	6.8	4.16
8b	9.9	3.5	34.0	-	-
9	3.9	7.0	35.7	5.3	1.27
10a	4.5	7.0	43.6	5.15	1.6
10b	11.1	3.7	33.0	1.00	0.3

30

Table 18

The Mg, Ti, DOP, EHA and PA contents of the catalysts in mol-% units					
Example	Mg mol-%	Ti mol-%	DOP mol-%	EHA mol-%	PA mol-%
8a	0.148	0.117	0.091	0.052	0.028
8b	0.407	0.073	0.087	-	-
9	0.161	0.146	0.091	0.041	0.009
10a	0.185	0.146	0.112	0.040	0.011
10b	0.457	0.077	0.085	0.008	0.002

45

Table 19

Comparison between the molar amounts of Mg, Ti and DOP			
Example	Mg/Ti	Ti	DOP/Ti
8a	1.3	1	0.8
8b	5.6	1	1.2
19	1.1	1	0.6

EP 0 926 165 A1

Table 19 (continued)

Comparison between the molar amounts of Mg, Ti and DOP			
Example	Mg/Ti	Ti	DOP/Ti
10a	1.3	1	0.8
10b	5.6	1	1.1

5

10

Table 20

The calculated amounts of Cl in the catalysts compared to the amounts found		
Example	Calculated w-%	Found w-%
8a	27.1	26.0
8b	39	-
9	32.1	30.9
10a	33.5	32.4
10b	43.4	44.0

15

20

25

#### The IR results

[0103] In Figure 9 is shown the IR spectra of the catalyst components of examples 9 and 10a. There were clear indications of the presence of phthalic anhydride in the catalyst that has been prepared from  $MgR_2$  (example 8a). The phthalic anhydride was almost totally missing from the samples that had been prepared out of  $MgCl_2$  (examples 9 and 10a). These results confirm the results of the chemical analysis. The IR spectrum (not shown) for the toluene washed example 10b catalyst showed no traces of phthalic anhydride but to the left of the  $C=O---Ti$  peak a shoulder had appeared indicating the presence of some free carboxylic acid group (-COOH).

35

#### The X-ray diffraction patterns

[0104] In Figures 10, 11 and 12, the X-ray diffraction patterns for the catalysts are shown. The results show that the addition of  $TiCl_4$  before PDC gives a more crystalline material. This can be seen in Figure 11. Example 9 is still showing the organic separation peak at  $7^\circ 2\Theta$  and the halo between  $18^\circ$  and  $22^\circ 2\Theta$  but only a slight remain thereof can be seen of the halo in the spectrum of example 10a. In all patterns there seems to be an additional peak at about  $32^\circ$ - $33^\circ 2\Theta$ . This peak is not connected to crystalline  $MgCl_2$ . Some unreacted  $MgCl_2$  seems to be present in the catalyst component of example 10a which is starting to dominate when the catalyst is washed with toluene (figure 12).

45

#### Polymerization results

[0105] All but one (example 8a) of the catalysts were test polymerized according to the descriptions in the experimental section. The polymerization results both in kg PP/g cat units and in kg PP/g Ti units are listed in Table 21. In Figure 13 the results are shown graphically. There was an almost logarithmic linear increase in the activities. As a whole it can be said that:

50

1. Addition of  $TiCl_4$  before PDC gives better activity (compare examples 9 and 10).
2. Starting from  $MgCl_2$  instead of from  $MgR_2$  gives higher activity (compare example 8 with examples 9 and 10).
3. Toluene wash improves activity (compare examples 10a and 10b).

Table 21

The polymerization results		
Example	Activity kg PP/g cat.	Activity kg PP/g Ti
8b	0.06	1.3
9	0.4	6.0
10a	1.2	18
10b	2.5	67

## Claims

15 1. A process for the preparation of an olefin polymerization catalyst component containing magnesium, titanium, halogen and an electron donor, **characterized by the steps of:**

20 (i) reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxide linked to magnesium, a complex containing a magnesium dihalide and an alcohol, and a non-complex magnesium dialkoxide, with a halogen compound (b) being capable of forming the electron donor by replacement of its halogen by said alkoxy moiety, to give an intermediate (ab), and

25 (ii) reacting said intermediate (ab) with a titanium halide (c), or

30 (i)' reacting a titaniumless magnesium compound (a) containing an alkoxy moiety, which titaniumless magnesium compound is selected from the group consisting of a compound or complex containing halogen and alkoxide linked to magnesium, and a complex containing a magnesium dihalide and an alcohol, with a titanium halide (c), to give an intermediate (ac), and

35 (ii)' reacting said intermediate (ac) with a halogen compound (b) being capable of forming the electron donor by replacement of its halogen by said alkoxy moiety.

40 2. A process according to claim 1, **characterized in that** at least one, preferably all of steps (i), (ii), (i)' and (ii)', are carried out in solution.

45 3. A process according to claim 1 or 2, **characterized in that** the reaction product of step (ii) or step (ii)' is recovered by precipitation.

50 4. A process according to claim 1, 2 or 3, **characterized in that** said compounds (a), (b) and (c) are contacted in essentially stoichiometric amounts, or alternatively, that a stoichiometric excess, preferably a 5-20 fold stoichiometric excess with respect to said titaniumless magnesium compound (a), of said titanium halide (c) is used.

55 5. A process according to any of claims 1-4, **characterized in that** said magnesium dihalide is magnesium dichloride  $MgCl_2$ .

6. A process according to any of claims 1-5, **characterized in that** said halogen compound (b) is an organic acid halide and preferably phthalic acid dichloride  $Ph(COCl)_2$ , wherein  $Ph$  is o-phenylene, whereby the electron donors formed therefrom are, correspondingly, an organic acid ester and preferably a phthalic acid diester  $Ph(COOR)_2$ , wherein R is a  $C_1-C_{20}$  alkyl or a  $C_7-C_{27}$  aralkyl, more preferably a di- $C_6-C_{16}$  alkyl phthalate, most preferably dioctyl phthalate.

7. A process according to any of claims 1-6, **characterized in that** said titanium halide (c) is a titanium tetrahalide, preferably titanium tetrachloride  $TiCl_4$ .

8. A process according to any of claims 1-7, **characterized by the steps of:**

(i) reacting said compound or complex containing halogen and alkoxide linked to magnesium as said titaniumless magnesium compound (a) with said halogen compound (b) to give an intermediate (ab) and

(ii) reacting said intermediate (ab) with said titanium halide (c), or:

5 (i)' reacting said complex of said magnesium dihalide and said magnesium dialkoxide as said titaniumless magnesium compound (a) with said titanium halide (c) to give an intermediate (ac) and  
(ii)' reacting said intermediate (ac) with said halogen compound (b).

9. A process according to claim 8, characterized in that said compound or complex containing halogen and alkoxide linked to magnesium as said titaniumless magnesium compound (a) is a complex of a magnesium dihalide and a magnesium dialkoxide, preferably a magnesium dichloride-dimagnesium dialkoxide complex  $MgCl_2 \cdot [Mg(OR)_2]_2$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, most preferably a  $C_6$ - $C_{16}$  alkyl, which complex is preferably prepared by reacting magnesium dichloride  $MgCl_2$  with an alcohol ROH to give an intermediate and reacting the obtained intermediate with a dialkyl magnesium  $MgR'''_2$ , wherein  $R'''$  is defined as for R.

10. A process according to claim 9, characterized by:

15 (i)' reacting said magnesium dichloride-dimagnesium dialkoxide complex  $MgCl_2 \cdot [Mg(OR)_2]_2$  (a) wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, with said titanium halide (c), which is said titanium tetrachloride  $TiCl_4$ , to give an intermediate (ac) and  
20 (ii)' reacting said intermediate (ac) with said halogen compound (b), which is said phthalic acid dichloride  $Ph(COCl)_2$ , wherein Ph is o-phenylene.

11. A process according to any of claims 1-7, characterized by the steps of:

25 (i) reacting said titaniumless magnesium compound (a) which is selected from said complex of said magnesium dihalide and said alcohol, and said non-complex magnesium dialkoxide, with said halogen compound (b) to give an intermediate (ab) which is a complex of said magnesium dihalide and said electron donor and  
(ii) reacting said intermediate (ab) which is a complex of said magnesium dihalide and said electron donor with said titanium halide (c).

30 12. A process according to claim 11, characterized in that, independently, said complex of said magnesium dihalide and said alcohol is a magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_m$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, and m is 1-6, and said non-complex magnesium alkoxide is a magnesium dialkoxide  $Mg(OR)_2$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, which is preferably prepared by reacting a magnesium dialkyl and an alcohol ROH.

35 13. A process according to any of claims 1-7, characterized by the steps of:

40 (i)' reacting said titaniumless magnesium compound (a), which is said complex of said magnesium dihalide and said alcohol, with said titanium halide (c) to give an intermediate (ac) and  
(ii)' reacting said intermediate with said halogen compound (b).

45 14. A process according to claim 13, characterized in that said complex of said magnesium dihalide and said alcohol compound is a magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_m$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, and m is 1-6.

15. A process according to claim 14, characterized in by the steps of:

50 (i)' reacting said titaniumless magnesium compound (a), which is said magnesium dichloride-alcohol complex  $MgCl_2 \cdot (ROH)_m$ , wherein R is a  $C_1$ - $C_{20}$  alkyl or a  $C_7$ - $C_{27}$  aralkyl, preferably a  $C_6$ - $C_{16}$  alkyl, and m is 1-6, with said titanium dihalide (c), which is said titanium tetrachloride  $TiCl_4$ , to give an intermediate (ac) and  
(ii)' reacting said intermediate (ac) with said halogen compound (b), which is said phthalic acid dichloride  $Ph(COCl)_2$ , wherein Ph is o-phenylene.

55 16. A process according to any of claims 8-15, characterized in that in step (ii) said intermediate (ab) is added, preferably drop by drop, to said titanium halide (c), which is in liquid form and preferably hot, most preferably at 75-150 °C.

17. A process according to any of claims 8-16, characterized in that:

EP 0 926 165 A1

(iii) the obtained reaction product of step (ii) or (ii)' is further treated with said titanium halide (c) and/or washed, preferably repeatedly washed with an aromatic hydrocarbon, such as toluene, or an organic liquid having the same solubility parameter as said aromatic hydrocarbon.

5 18. A catalyst component comprising magnesium, titanium, a halogen and an electron donor, characterized in that it has been prepared by the process described in any of claims 1 to 17.

10 19. The catalyst component according to claim 18, characterized in that it is an isolated complex of a magnesium dihalide, preferably magnesium dichloride, an electron donor obtained by replacing the halogen of a halogen compound by an alkoxy group, preferably a phthalic acid diester, and a titanium halide, preferably titanium tetrachloride.

15 20. The catalyst component according to claim 19, characterized in that said complex has an X-ray pattern comprising a crystal height indicating peak between 16° and 18° 2Θ (Siemens D500 instrument, CuK $\alpha$  radiation wavelength 1.541 Å, effect 40 kV and 35 mA).

20 21. The catalyst component according to claim 18, 19 or 20, characterized in that said complex is prepared by contacting stoichiometric amounts of said components (a), (b) and (c).

25 22. The catalyst component according to any of claims 18-21, characterized in that said complex has an X-ray diffraction pattern comprising a peak between 5° and 10° 2Θ (Siemens D500 instrument, CuK $\alpha$  radiation wavelength 1.541 Å, effect 40 kV and 35 mA).

25 23. The catalyst component according to any of claims 18-22, characterized in that no TiCl<sub>3</sub>OR waste material is produced in the catalyst synthesis.

30 24. Use of a catalyst component according to any of claims 18-23 for the polymerization of  $\alpha$ -olefins, preferably propene.

25 25. Use according to claim 24, wherein, additionally, a catalyst component comprising an organometal compound of a metal belonging to Group 1, 2 or 13 (IUPAC 1990) of the Periodic Table, preferably an alkyl aluminium compound, is used.

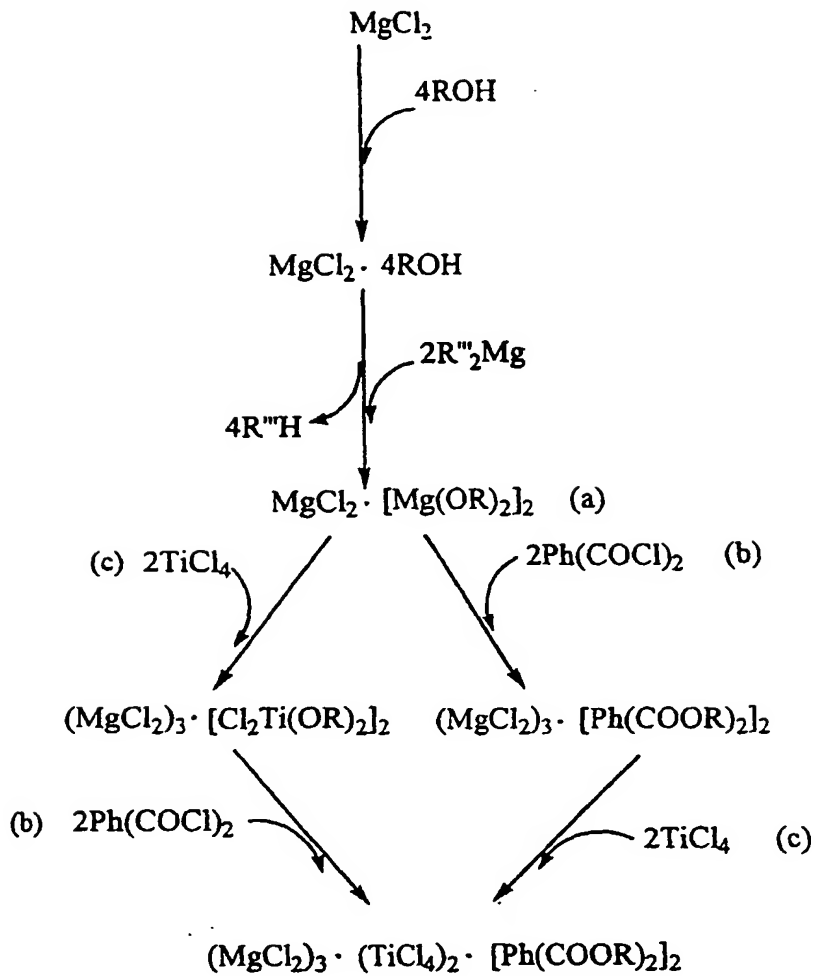
35

40

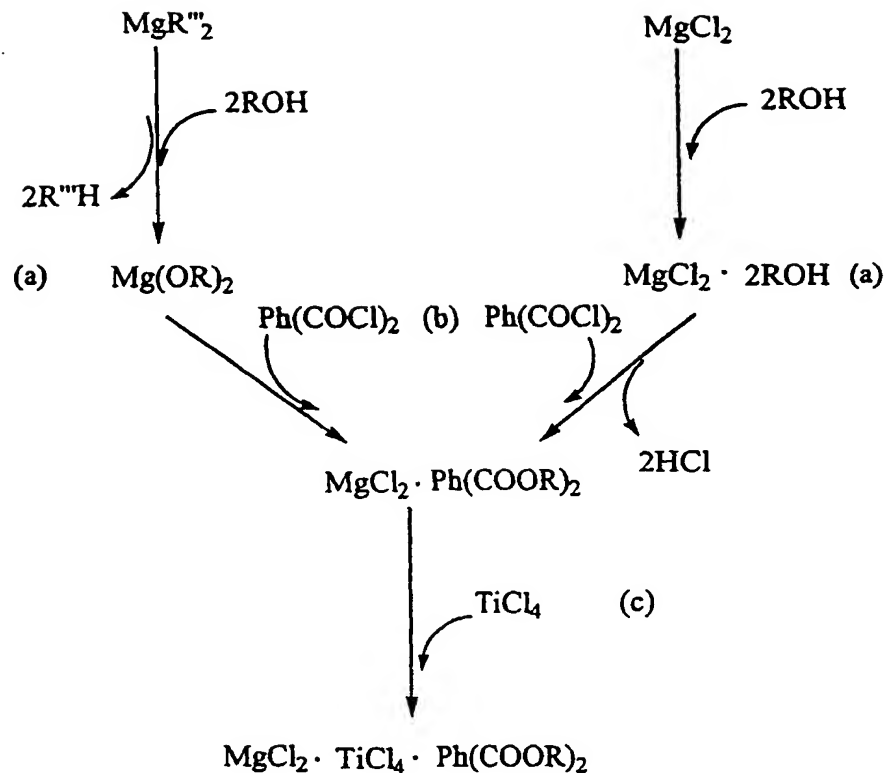
45

50

55



**Figure 1** Example of the first main embodiment



**Figure 2** Example of the second main embodiment

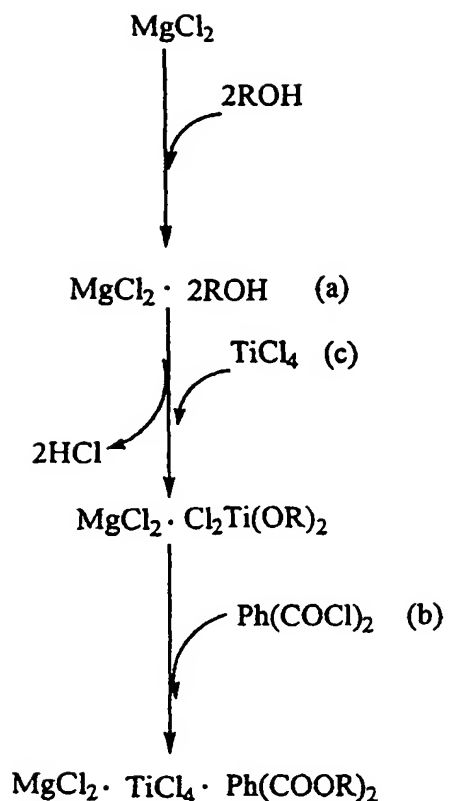
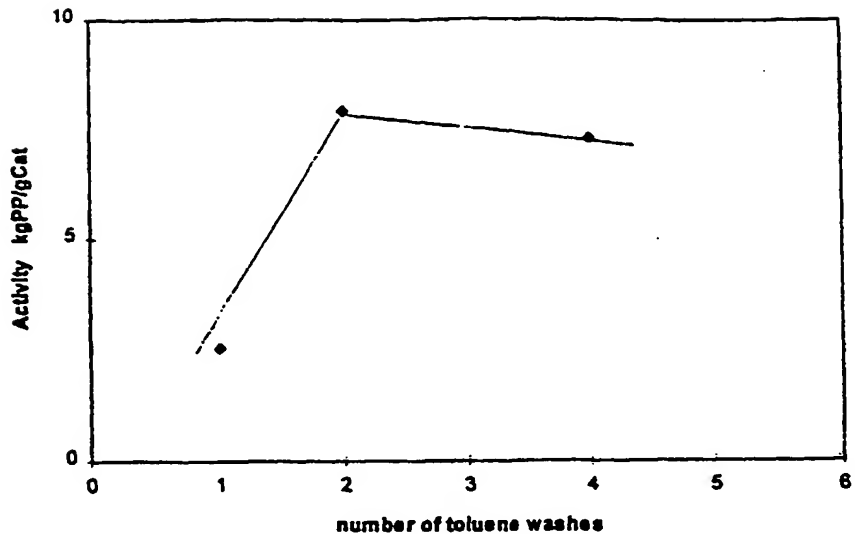
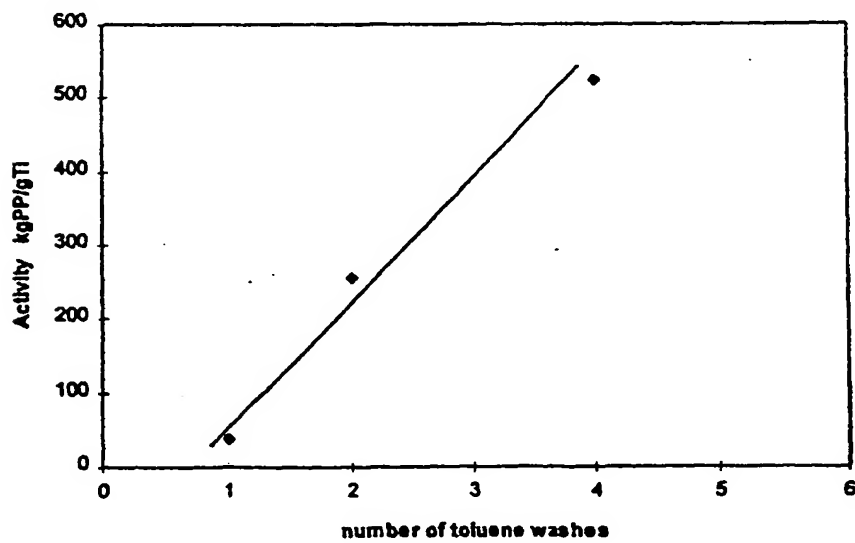


Figure 3 Example of the third main embodiment



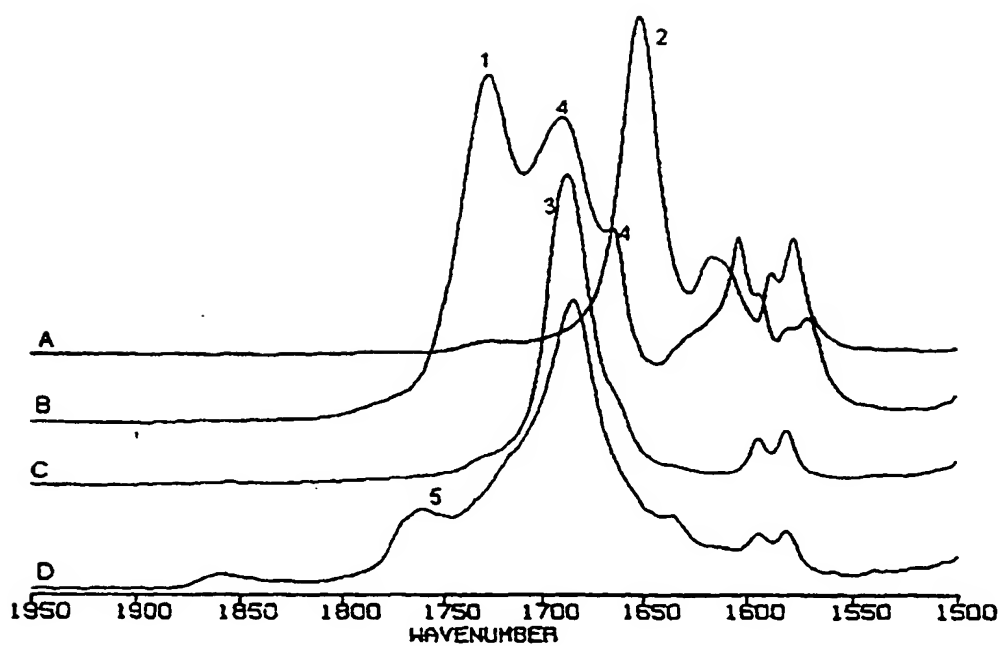
**Figure 4**

Correlation between the activity (in kg PP/g cat) and the number of toluene washes used in the catalyst synthesis



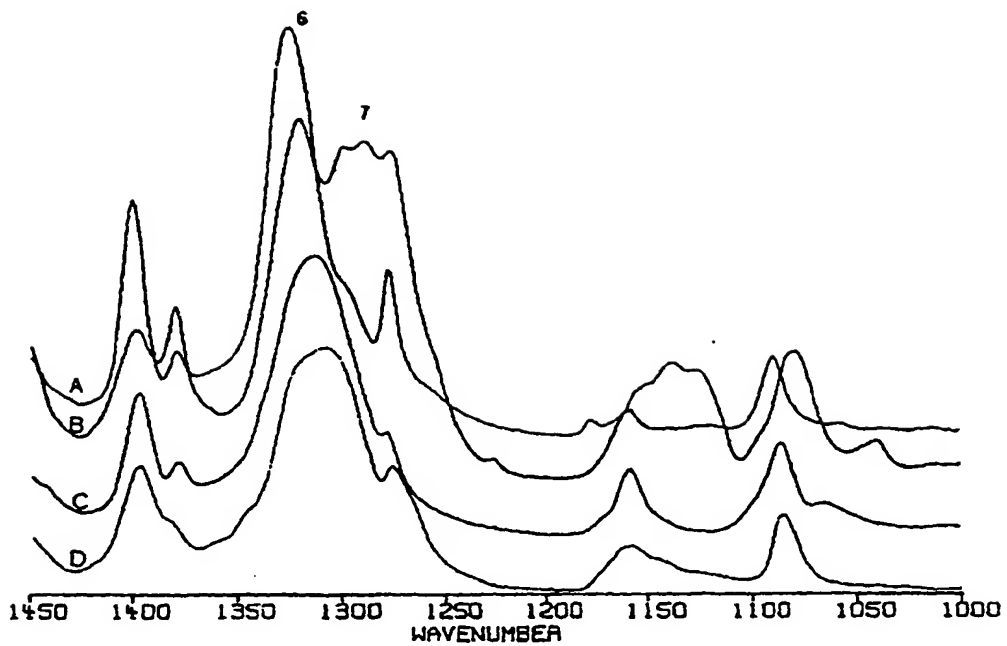
**Figure 5**

Correlation between the activity (in kg PP/g Ti) and the number of toluene washes used in the catalyst synthesis



**Figure 6**

Details of IR spectra in the C=O—metal complex region of  $\text{TiCl}_4 \cdot \text{DUP}$  (A),  $(\text{MgCl}_2)_{1.5} \cdot \text{DOP}$  (B),  $\text{MgCl}_2 \cdot \text{TiCl}_4 \cdot \text{D}_1$  (C) prepared from  $\text{MgCl}_2 \cdot 3\text{ROH}$ , and of  $(\text{MgCl}_2)_7 \cdot \text{TiCl}_4 \cdot \text{DOP}$  (D = example 6) prepared from  $\text{MgCl}_2 \cdot \text{DOP}$  in 10 mol  $\text{TiCl}_4$



**Figure 7**

Details of IR spectra in the C-O--metal complex region of  $\text{TiCl}_4 \cdot \text{DUP}$  (A),  $(\text{MgCl}_2)_{1.5} \cdot \text{DOP}$  (B),  $\text{MgCl}_2 \cdot \text{TiCl}_4 \cdot \text{D}_1$  (C) prepared from  $\text{MgCl}_2 \cdot 3\text{ROH}$ , and of  $(\text{MgCl}_2)_7 \cdot \text{TiCl}_4 \cdot \text{DOP}$  (D = example 6) prepared from  $\text{MgCl}_2 \cdot \text{DOP}$  and 10  $\text{TiCl}_4$

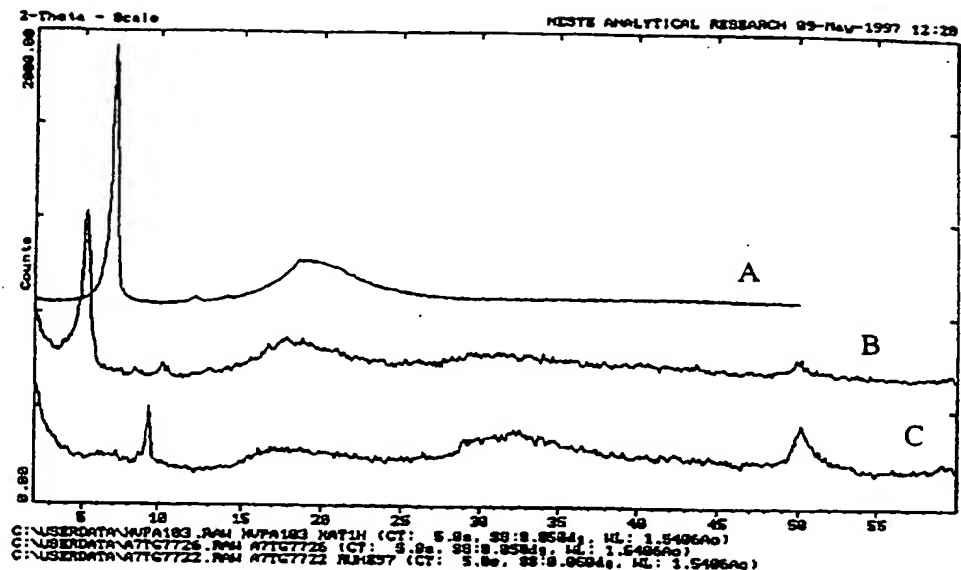


Figure 8

X-ray patterns of  $\text{Mg}(\text{OR})_2$  (A),  $\text{MgCl}_2 \cdot \text{TiCl}_4 \cdot \text{DOP}$  (B) produced from  $\text{MgCl}_2 \cdot 3\text{EtOH}$ , and  $(\text{MgCl}_2)_{1.7} \cdot \text{TiCl}_4 \cdot \text{DOP}$  (C) produced from  $\text{MgCl}_2 \cdot \text{DOP}$  and 10  $\text{TiCl}_4$

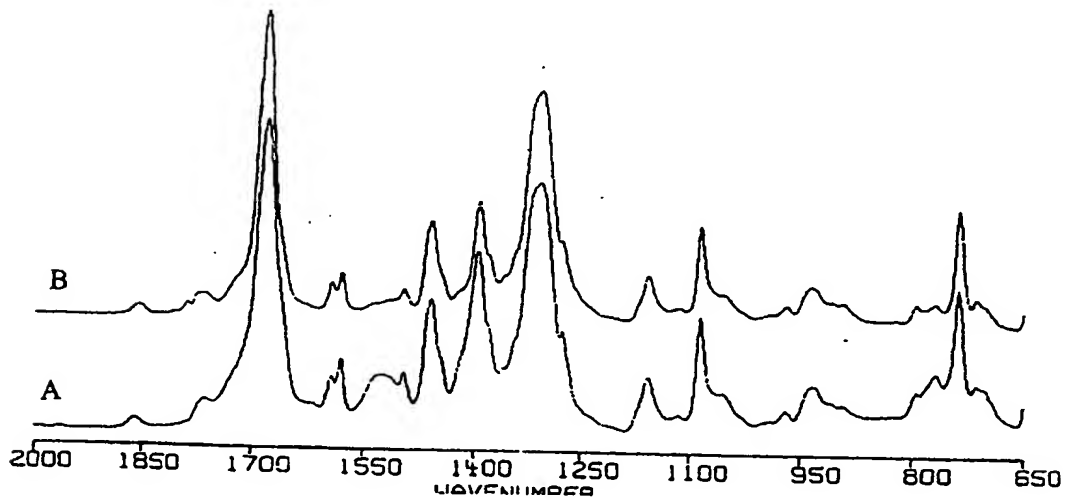


Figure 9

IR spectra of examples 9 (A) and 10a (B)

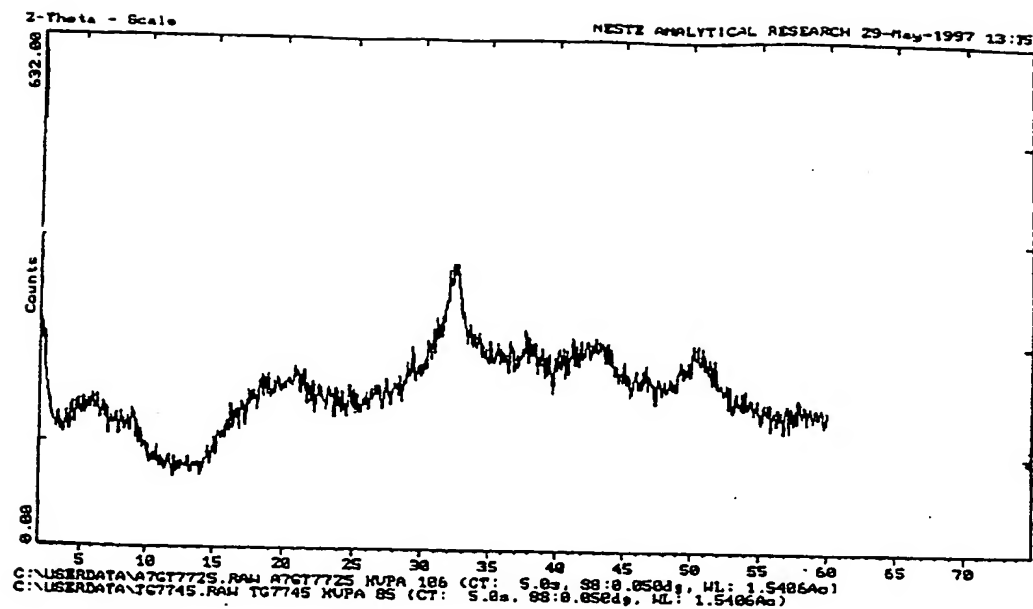


Figure 10

X-ray pattern of the catalyst of example 8a

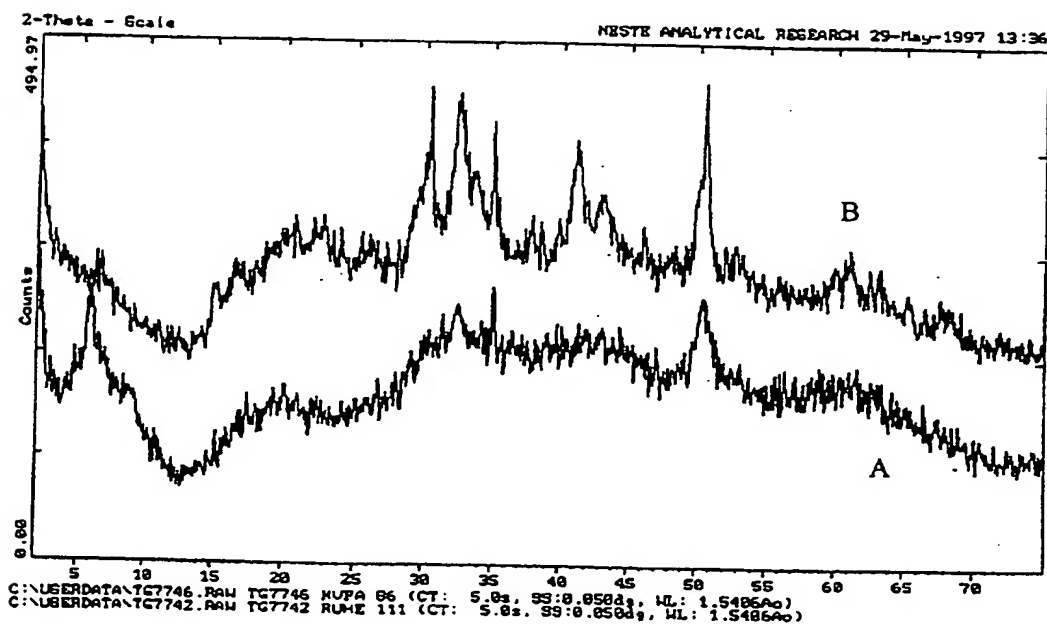


Figure 11

X-ray patterns of examples 9 (A) and 10a (B)

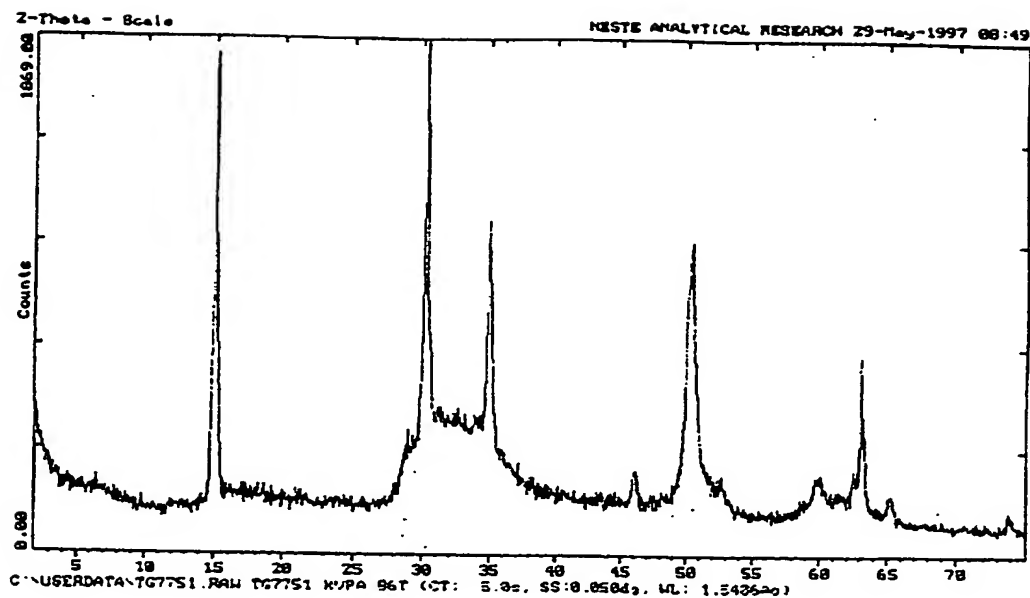


Figure 12

X-ray pattern of example 10b, i.e. the toluene washed example 10a

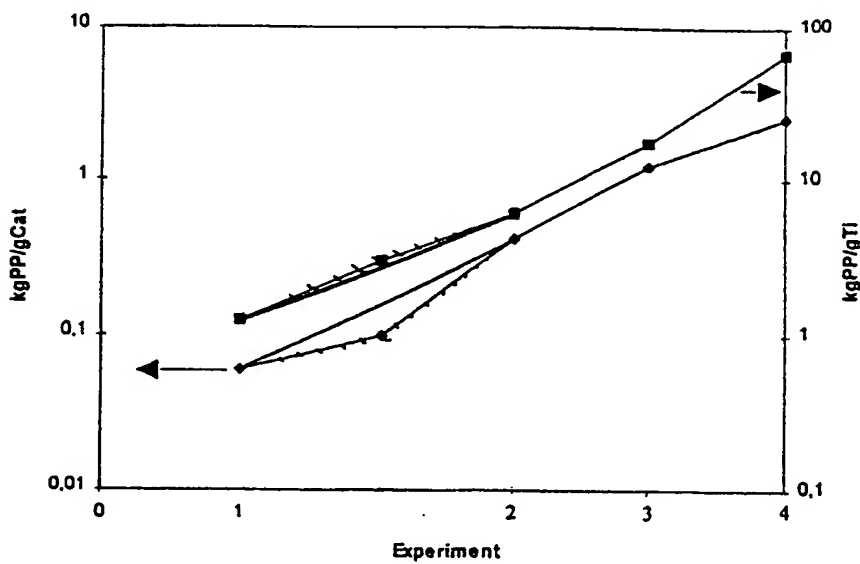


Figure 13

The activities of the catalysts coming from experiments 8b-10b,  
 1) example 8b, 2) example 9, 3) example 10a and 4) example 10b.  
 ◆ as kg PP/g cat units and ■ as kg PP/g Ti units



European Patent  
Office

## EUROPEAN SEARCH REPORT

Application Number

EP 98 66 0147

DOCUMENTS CONSIDERED TO BE RELEVANT			CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	
X	EP 0 195 497 A (TOA NENRYO KOGYO KK) 24 September 1986  * example 27 * * examples 1,2 * * claims 1-4,6,10 * ---	1,6,7, 11,12, 17-19, 24,25	C08F4/654 C08F10/00
X	EP 0 288 762 A (BASF AG) 2 November 1988  * example 1 * ---	1,5-7, 11,13, 17-19, 24,25	
X	EP 0 743 326 A (MITSUI PETROCHEMICAL IND) 20 November 1996  * comparative example 6 * * example 12 * * page 3, line 10 - line 11 * * table 5 * ---	1,2,5-7, 13-15, 18,24,25	
A	EP 0 171 200 A (TOA NENRYO KOGYO KK) 12 February 1986 * example 46 * -----	1	C08F
The present search report has been drawn up for all claims			TECHNICAL FIELDS SEARCHED (Int.Cl.6)
Place of search	Date of completion of the search	Examiner	
THE HAGUE	25 March 1999	Fischer, B	
CATEGORY OF CITED DOCUMENTS		T : theory or principle underlying the invention E : earlier patent document, but published on or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document	
X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document			

EP 0 926 165 A1

ANNEX TO THE EUROPEAN SEARCH REPORT  
ON EUROPEAN PATENT APPLICATION NO.

EP 98 66 0147

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on. The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

25-03-1999

Patent document cited in search report	Publication date	Patent family member(s)			Publication date
EP 0195497 A	24-09-1986	JP	1973613 C		27-09-1995
		JP	7002775 B		18-01-1995
		JP	61174204 A		05-08-1986
		AU	611846 B		27-06-1991
		AU	5276186 A		31-07-1986
		CA	1263361 A		28-11-1989
EP 0288762 A	02-11-1988	DE	3711919 A		27-10-1988
		AT	87011 T		15-04-1993
		AU	1436988 A		13-10-1988
		CA	1302639 A		02-06-1992
		DE	3879257 A		22-04-1993
		ES	2053600 T		01-08-1994
		JP	63265903 A		02-11-1988
		US	4843132 A		27-06-1989
EP 0743326 A	20-11-1996	CA	2176896 A		19-11-1996
		CN	1141304 A		29-01-1997
		JP	9031119 A		04-02-1997
EP 0171200 A	12-02-1986	JP	1927368 C		25-04-1995
		JP	6055780 B		27-07-1994
		JP	61021109 A		29-01-1986
		AU	582169 B		16-03-1989
		AU	4467985 A		16-01-1986
		CA	1248511 A		10-01-1989
		US	4727049 A		23-02-1988